

Inhibitors of the leukocyte proMMP-9/beta(2) integrin complex

Field of the Invention

5 The present invention concerns peptide compounds, which are inhibitors of integrin-MMP complex. In specific, the compounds bind to the α_M integrin I domain and inhibit its complex formation with proMMP-9. The compounds thus prevent neutrophil migration, as well as leukocyte migration. The compounds can be used in the treatment of inflammatory conditions, and leukaemia.

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Background of the Invention

The leukocyte integrin family consists of four heterodimeric glycoproteins with specific α -chains (α_L , α_M , α_X , or α_D) and a common β_2 -chain (CD18). They play an 15 essential role in mediating adhesion of cells in the immune system (1). The major ligand-binding site locates to an 200 amino acid long sequence within the α -chain called I or inserted domain, which is homologous to the A domains of von Willebrand factor, repeats of cartilage matrix protein and collagen (2).

20 Among the β_2 integrins, $\alpha_M\beta_2$ is the most promiscuous binder being able to interact with a multitude of unrelated ligands. These include ICAM 1 to 5, complement fragment iC3b, fibrinogen, uPAR, E-selectin and various extracellular matrix proteins (see (3) and references therein). The integrin has also been shown to have a capacity to bind certain enzymes, but whether this is important for leukocyte adhesion or immune 25 reactions is unclear. Such enzymes showing integrin-binding activity are catalase (4), myeloperoxidase (5) and the proteinases elastase (6) and urokinase (7).

Extensive work has been done to identify the ligand binding sites in β_2 integrin I-domains, but less is known about the interacting ligand regions (8, 9). Recently, the 30 structure of an α_L I domain/ICAM-1 complex was reported (10). Low molecular weight peptides binding to the β_2 integrins are useful reagents to study integrin function, and such peptides have been derived from ICAM-2 (11), fibrinogen (12), and Cyr61 (13). We have used phage display libraries to study the peptide binding specificity of

integrins and to develop potential drug leads. In our previous study, we isolated the bicyclic peptide CPCFLLGCC (SEQ ID NO: 1) (LLG-C4)(unpublished observations of the present inventors) as the most active binder to the purified $\alpha_M\beta_2$ integrin (14). Leukocytes can efficiently adhere to the immobilized LLG-C4 peptide via the $\alpha_M\beta_2$ and

5 $\alpha_X\beta_2$ integrins.

Summary of the Invention

We have now extended phage display screenings to the purified α_M I domain. This has
10 resulted in the identification of a novel I domain-binding tetrapeptide motif D/E-D/E-G/L-W (SEQ ID NO: 2), which is found on some of the known β_2 integrin ligands and interestingly also on the catalytic domain of MMPs. We show that the D/E-D/E-G/L-W (SEQ ID NO: 2) motif mediates binding between an MMP and β_2 integrin, and proMMP-9 gelatinase, which is known to be the major MMP of leukocytes, occurs in
15 complex with the $\alpha_M\beta_2$ and $\alpha_L\beta_2$ integrins in leukaemic cell lines following cellular activation. The peptide inhibitors of the integrin-MMP complex prevent leukaemia cell migration, suggesting a role for the complex in cell motility. The compounds of the invention also attenuated PMN migration *in vitro* and *in vivo*, suggesting a role for the MMP-integrin complex in PMN motility.

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Detailed description of the Invention

Analysis of α_M I domain-binding peptides led to the finding that MMPs, particularly the
25 MMP-9 and MMP-2 progelatinases, are potent β_2 integrin ligands. Our studies show that proMMP-9, the major MMP of activated leukocytes, is co-localized with the β_2 integrin on the cell surface. Cell surface labelling and co-immunoprecipitation further demonstrates the occurrence of the complex in leukemic cell lines. Finally, we have found evidence that this proteinase-integrin complex plays a role in migration of the leukemic cells.

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Although phage display has been extensively used with whole integrins, to the best of our knowledge, this is the first successful phage display selection on an isolated integrin I domain. We could enrich only one binding motif even though the α_M I domain can

- bind a variety of ligands. The peptide motif we isolated could not compete with the ICAM-1, fibrinogen, or LLG-C4 ligands. The success of phage display depends on the libraries used and the biopanning conditions. Our method favoured “high affinity” interactions with the cyclic peptides yielding the D/E-D/E-G/L-W (SEQ ID NO: 2) motif. Interestingly, this motif shows a high degree of similarity to the CWDD(G/L)WLC (SEQ ID NO: 3) peptide isolated by phage display as an RGD sequence-binding peptide (15). By recognizing the RGD ligand sequence, CWDDGWLC (SEQ ID NO: 3) structurally and functionally behaves like a minimal integrin. Here, we have identified the DDGW (SEQ ID NO: 174) peptide in a reverse situation, as a ligand to integrin. However, the RGD sequence does not compete with the α_M I domain as the GRGDSP (SEQ ID NO: 4) peptide at a 1 mM concentration was unable to inhibit proMMP-9 binding to the I domain (unpublished observations of the present inventors).
- 15 Some hints for the binding site of DDGW (SEQ ID NO: 174) come from the interaction of iC3b with $\alpha_M\beta_2$ integrin. Our pepspot analysis showed that the iC3b peptide ARSNLDEDIIAEENI (SEQ ID NO: 5), but not the control peptide ARSNLDAAIIAEENI (SEQ ID NO: 6), bound the α_M I domain and the DDGW (SEQ ID NO: 174) peptide blocked this binding. The DEDIIAEENI (SEQ ID NO: 7) sequence with multiple adjacent negative charges is required for efficient binding of iC3b to $\alpha_M\beta_2$ integrin. The binding site of complement protein iC3b in the I domain has been mapped indicating a role for the positively charged amino acid residue K²⁴⁵ for iC3b binding. Mutation of this residue does not affect the binding of the fibrinogen recognition peptide. This may account for the inability of the DDGW (SEQ ID NO: 174) peptide to inhibit ICAM-1 and fibrinogen-mediated cell adhesion. These findings suggest that the K²⁴⁵ residue is the positively charged contact site for the DDGW (SEQ ID NO: 174) peptide and the D/E-D/E-G/L-W (SEQ ID NO: 2) motif as well.

The pepspot analysis indicates that a class of β_2 integrin ligands contains an active D/E-D/E-G/L-W (SEQ ID NO: 2) motif. These include the previously identified $\alpha_M\beta_2$ ligands iC3b, thrombospondin-1, and the enzymes myeloperoxidase and catalase. In our experiments, the peptides derived from several secreted MMPs, but not membrane-bound MT1-MMP, were also active. It is notable that the D/E-D/E-G/L-W (SEQ ID

NO: 2) motif is relatively conserved in the secreted members of the MMP family.

Finding of a dominant integrin-binding site in the catalytic domain of proMMP-9 was unexpected, because previous studies suggested an essential role for another MMP 5 domain, the hemopexin domain, in integrin binding. The hemopexin domains mediated MMP-2 binding to the $\alpha_v\beta_3$ integrin and MMP-1 binding to the $\alpha_2\beta_1$ integrin. The cleaved hemopexin domain of MMP-2 has also been shown to occur *in vivo* and to inhibit angiogenesis. Understandably, phage peptide display and pepspot techniques have limitations and only linear peptide sequences can be analysed, not protein 10 conformations. Thus, in the present study we cannot make conclusions of the function of separate MMP-9 domains in integrin binding and it remains to be seen whether cleavage products of MMP-9, if present *in vivo*, can act as β_2 integrin ligands. Our studies suggest that the peptide sequence from the catalytic domain is essential for the 15 binding of full-length proMMP-9 to the β_2 integrin, as the synthetic DDGW (SEQ ID NO: 174) peptide could completely inhibit the integrin binding. It was important to use natural proMMP-9 because $\alpha_M\beta_2$ is known to bind to denatured proteins and this may sometimes be the case for bacterially expressed proteins. Furthermore, we did not observe binding of an active MMP-2 or MMP-9 to the integrin, although the DELW(T/S)LG (SEQ ID NO: 8) sequence should remain unchanged in the active 20 enzyme. We rather found that AMPA and trypsin, activators of proMMP-9, released MMP-9 from THP-1 cells, apparently affecting the integrin complex (unpublished observations of the present inventors). These results suggest that the proenzyme presents the integrin binding site more efficiently than the active enzyme and the β_2 integrin may even control the activation of the proenzyme.

25 In the three-dimensional structures of proMMP-2 and -9, the I domain binding site is located in the vicinity of the zinc-binding catalytic sequence HEFGHALGLDH (SEQ ID NO: 9) between the catalytic domain and the fibronectin type II repeats. This location suggests a mechanism for evading proMMP-9 inhibition by tissue inhibitors of 30 MMPs (TIMPs) or α_2 -macroglobulin. In the absence of inhibitors, the cell surface-localized proMMP-9 would be readily susceptible for activation and substrate hydrolysis, which may also occur in the presence of intact propeptide. On the other hand, because the binding site of the I domain is located in the vicinity of the catalytic

groove, it also suggests an explanation for the blocking of MMP-9/β₂ integrin interaction by the small molecule MMP inhibitors such as CTT and Inh1.

- The activity of the DDGW (SEQ ID NO: 174) peptide in the THP-1 cell migration assay suggests an important function for the integrin-progelatinase complex in leukocyte migration. Obviously, we cannot exclude the possibility that the DDGW (SEQ ID NO: 174) peptide blocks binding of other ligands than gelatinases and in this way inhibits the leukocyte migration. However, as the specific gelatinase inhibitor CTT also blocks the THP-1 cell migration, these results strongly suggest that the proMMP-9/β₂ integrin complex is the main target for DDGW (SEQ ID NO: 174). Interestingly, the DDGW (SEQ ID NO: 174) peptide blocked THP-1 cell migration although it increased the level of proMMP-9 in the medium, which suggests that cell-surface bound rather than total MMP-9 level is a critical factor in cell migration.
- DDGW (SEQ ID NO: 174) and HFDDDE (SEQ ID NO: 10) (see below) had potent activities *in vivo* in the mouse peritonitis model, but it is unclear to what extent this was due to inhibition of proMMP-9 as both peptides can potentially inhibit other β₂ integrin ligands as well. A subset of β₂ integrin ligands have a DDGW(SEQ ID NO: 174) like sequence and these include, in addition to MMPs, at least complement iC3b and thrombospondin-1. The excellent *in vivo* activity of DDGW (SEQ ID NO: 174) makes it a useful tool to study the components involved in leukocyte migration and the peptide may be considered as a lead to develop anti-inflammatory compounds. Our results suggest that the proMMP-9/α_Mβ₂ complex may be part of the neutrophil's machinery for a specific β₂ integrin-directed movement.
- The present invention is thus directed to new peptide compounds, in specific to peptide compounds comprising the tetrapeptide motif D/E-D/E-G/L-W (SEQ ID NO: 2). Said compounds can be used as pharmaceuticals, which inhibit leukocyte migration, as well as neutrophil migration. The inhibitory activity was shown both in *in vitro* and *in vivo* experiments. Consequently, the compounds can be used to treat leukaemia and prevent and treat inflammatory conditions.

One embodiment of the invention is the use of the compounds of the invention for the manufacture of a pharmaceutical composition for prophylaxis and treatment of

conditions dependent on neutrophil migration.

Another embodiment of the invention is the use of the compounds of the invention for the manufacture of a pharmaceutical composition for the treatment of conditions
5 dependent on leukocyte migration.

A further embodiment of the invention is a pharmaceutical composition comprising as an active ingredient a compound of the invention, and a pharmaceutically acceptable carrier.

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A still further embodiment of the invention is a method for therapeutic or prophylactic treatment of conditions dependent on leukocyte or neutrophil migration, comprising administering to a mammal in need of such treatment a leukocyte or neutrophil migration inhibiting compound of the invention in an amount which is effective in
15 inhibiting migration of leukocytes or neutrophils. Specific embodiments of the invention include methods for treatment of leukaemia and inflammatory conditions.

Abbreviations used

- 20 APMA, aminophenylmercuric acetate;
 $\alpha_M\beta_2$, CD11b/CD18, Mac-1 integrin;
CTT, CTTHWGFTLC (SEQ ID NO: 11) peptide;
DDGW(SEQ ID NO: 174), ADGACILWMDDGWCAGAAG(SEQ ID NO: 12) peptide;
GPA, glycolphorin A;
25 GST, glutathione-S-transferase;
HMEC, human microvascular endothelial cell;
ICAM, intercellular adhesion molecule;
Inh1, matrix metalloproteinase inhibitor 1;
KKGW(SEQ ID NO: 13), ADGACILWMKKWCGAAG (SEQ ID NO: 14)peptide;
30 LLG-C4, CPCFLLGCC (SEQ ID NO: 1)peptide;
MMP, matrix metalloproteinase;
NGAL, neutrophil gelatinase-associated lipocalin;
PMN, polymorphonuclear neutrophil;
RGD-C4, ACDCRGDCFCG (SEQ ID NO: 15)_peptide;

STT, STTHWGFTLS (SEQ ID NO: 16)_peptide;
 TAT-2: tumor-associated trypsinogen-2;
 W→A CTT, CTTHAGFTLC (SEQ ID NO: 17)_peptide.

5 Description of the Drawings

FIG. 1A to 1E. Identification of an I domain binding site in progelatinase.

FIG. 1A, Phage display peptide sequences (SEQ ID NOS: 105-108, 20, 109-110, 19, 111-115, 20, 19, 116-123 respectively) specifically bound to the α_M I domain. The consensus motif is shown in bold. Peptides with the strongest binding (CILWMDDGW) (SEQ ID NO: 18) and the highest similarity (CPEELWWLC) (SEQ ID NO: 19) are aligned with human MMPs (accession numbers shown in parenthesis).

FIG. 1B, Phages bearing the CILWMDDGWC (SEQ ID NO: 20) peptide or a control peptide were allowed to bind to immobilized α_M I domain-GST fusion protein (20 ng/well) in the absence or presence of 15 μ M DDGW (SEQ ID NO: 174)_peptide or LLG-C4 peptide. Bound phages were detected using a monoclonal anti-M13 phage antibody. Mean absorbance of triplicate samples \pm SD is shown.

FIG. 1C, α_L , α_M , or α_X I domain-GST fusions were coated on microtiter wells as in B, and the binding of CILWMDDGWC (SEQ ID NO: 20)_peptide bearing phage or a control phage was measured.

FIG. 1D, Peptides (SEQ ID NOS: 124-167) covering the complete sequence of proMMP-9 were synthesized as overlapping peptides on a pepspot membrane. The α_M I domain (0.5 μ g/ml) was allowed to bind to the peptides followed by immunodetection using anti- α_M I domain antibody LM2/1. The α_M I domain-binding peptide 13 (arrow) is shown in boldface and the zinc binding catalytic sequence is underlined. The prodomain (Pro), catalytic domain containing the fibronectin type II repeats (Cat) and hemopexin domain (Pex) are marked to illustrate the domain structure of proMMP-9.

FIG. 1E, Alanine-mutated and truncated peptides (SEQ ID NOS: 2, 168-169, 20, 29, 170-173 respectively) were synthesized on a pepspot filter and probed with the recombinant α_M I domain (5 μ g/ml). Bound I domain was measured using mAb MEM-170 (5 μ g/ml) followed by HRP-conjugated anti-mouse secondary antibody and ECL detection. The binding was quantified by densitometric scanning. The bars show α_M I domain binding to single peptide spots as arbitrary optical density units/mm². Similar results were obtained in three independent experiments.

FIG. 2A to 2C. Binding of progelatinases to purified integrins and I domains.

FIG. 2A, proMMP-9, proMMP-2 or their trypsin-activated forms, at the concentrations indicated, were allowed to bind to $\alpha_M\beta_2$ integrin-coated wells. Appropriate MMP

5 antibodies were used to determine binding. The results in this and other figures are represented as the means \pm SD from triplicate wells.

FIG. 2B, binding of proMMP-9 (80 ng/well) was examined on microtiter wells coated with an integrin ($\alpha_L\beta_2$, $\alpha_M\beta_2$, $\alpha_1\beta_1$, $\alpha_3\beta_1$) or an I domain (α_L , α_M , α_X). The binding was determined using anti-MMP-9 antibody.

10 **FIG. 2C,** proMMP-2 (80 ng/well) was allowed to bind to $\alpha_L\beta_2$, $\alpha_M\beta_2$, $\alpha_1\beta_1$ or the I domains α_L , α_M or α_X . The binding was determined using an anti-MMP-2 antibody.

FIG. 3A to 3D. Inhibitors of the proMMP-9 / β_2 integrin complex.

FIG. 3A, α_M and α_L I domain-GST fusions were immobilized on microtiter wells.

15 ProMMP-9 (100 ng/well) was added in the presence or absence of various peptides (200 μ M) or lovastatin (100 μ M) in a buffer containing 0.5% BSA. ProMMP-9 binding was detected using a monoclonal antibody against MMP-9. The results are shown as percent binding compared to binding in the absence of inhibitors (100%) and no proMMP-9 added (0%).

20 **FIG. 3B,** DDGW (SEQ ID NO: 174) peptide blocks proMMP-9 binding to α_M in a dose dependent manner. The assay was done similarly as in (A), except various concentrations of peptides were added to compete for binding. All samples were assayed as triplicates and results shown are means \pm SD from a representative experiment.

25 **FIG. 3C,** proMMP-9 binding to $\alpha_M\beta_2$ and $\alpha_L\beta_2$ was examined in the absence and presence of EDTA (5 mM), MMP inhibitor-1 (100 μ M), CTT (200 μ M), STT (200 μ M), MEM170 (40 μ g/ml), or control TL3 antibody (40 μ g/ml). The background of primary and secondary antibodies was measured by omitting proMMP-9 from the wells or by coating with ICAM-1.

30 **FIG. 3D,** proMMP-9 binding to purified α_M and α_L I domain GST fusion proteins or wild type GST was studied in the absence or presence of competitors as indicated. Control shows the background when proMMP-9 was omitted.

FIG. 4A and 4B. Coprecipitation of progelatinase with β_2 integrin.

FIG. 4A, $\alpha_M\beta_2$ integrin (3 μ g) was incubated with a 500 μ l sample of HT1080 medium containing proMMP-9 and proMMP-2 in the absence or presence of CTT or STT (200 μ M) for 2 h. The integrin was immunoprecipitated with the OKM10 antibody, and the immunoprecipitates were analyzed by gelatin zymography. In control experiments, integrin was omitted from the medium and ICAM-1 was added instead.

FIG. 4B, a 500 μ l sample of HT1080 medium containing proMMP-9 and proMMP-2 was incubated with the α_M I domain GST (3 μ g) or LLG-C4-GST control. ICAM-1, LM2/1, CTT, STT, or LLG-C4 were used as competitors. GST was pulled down with glutathione beads, and bound proteins were analyzed by zymography. The lane 1 in the figure insert: the proMMP-2 and proMMP-9 zymogens present in non-treated HT1080 medium, lane 2: lack of gelatinases pulled down with control LLG-C4-GST, lane 3: proMMP-9 and proMMP-2 coprecipitated by α_M I domain GST fusion protein.

15 **FIG. 5A and 5B. CTT peptide binds to both latent and active MMP-9.**

FIG. 5A, Binding of proMMP-9 or APMA-activated MMP-9 to CTT-GST was examined in the absence or presence of competitors CTT (100 μ M), W \rightarrow A mutant CTT (100 μ M), and Inh1 (100 μ M). GST control was LLG-C4-GST. Binding was determined as in Figs. 2 and 3. The background in the absence of proMMP-9 is shown.

20 **FIG. 5B,** THP-1 cells were incubated in serum-free medium containing CTT, Inh1 or W \rightarrow A CTT at 200 μ M concentration. Samples from the media were collected at the time points indicated and analyzed by zymography (panels 1, 3, and 4) or Western blotting (panel 2).

25 **FIG. 6A to 6C. Progelatinases occur in complex with $\alpha_M\beta_2$ and $\alpha_L\beta_2$ in PDBu-activated THP-1 and Jurkat cells.**

FIG. 6A, THP-1 cell surface proteins were [3 H]-labelled using periodate-tritiated borohydride and analyzed by immunoprecipitation. CTT was used as a competitor (200 μ M). The immunoprecipitated samples were resolved on a 8-16% polyacrylamide gel, 30 and the film was exposed for 3 days. Lanes 1-4 are from non-activated cells and lanes 6-10 from PDBu-activated cells. Lane 5 shows molecular weight markers.

FIG. 6B, lysates from PDBu-activated THP-1 cells were immunoprecipitated with integrin or MMP antibodies followed by Western blotting with α_M (OKM10), α_L

(TS2/4) or MMP-9 antibodies. Preclearings of the cell lysates were done using α_M (lane 6) and α_L (lane 7) antibodies.

FIG. 6C, lysates from PDBu-activated Jurkat cells were subjected to immunoprecipitation followed by blotting with the α_L (MEM83) and MMP-9 antibodies.

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FIG. 7A and 7B. PDBu-induced colocalization of $\alpha_M\beta_2$ and proMMP-9 in THP-1 cells. Cells were preincubated for 30 min at +37°C with 50 nM PDBu.

FIG. 7A, cells were treated with anti- α_M OKM10 and anti-MMP-9 antibodies followed by FITC-labeled (green fluorescence) and TRITC-labeled (red fluorescence) secondary 10 antibodies. Yellow color indicates colocalization of $\alpha_M\beta_2$ and proMMP-9. Bars, 8.5 μ m.

FIG. 7B, immunofluorescence staining shows intense colocalization of MMP-9 (polyclonal antibody) and $\alpha_M\beta_2$ integrin (OKM-10) on the surface of PDBu-activated THP-1 cells at higher magnification as visualized by confocal microscopy (Bars, 2.5 μ m).

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FIG. 8A to 8D. The DDGW (SEQ ID NO: 174) peptide supports THP-1 cell adhesion and induces proMMP-9 release, but does not block adhesion to the major β_2 integrin ligands fibrinogen and ICAM-1. KKGW is SEQ ID NO: 13.

FIG. 8A, THP-1 cells were allowed to bind to immobilized, glutaraldehyde 20 polymerized peptides with or without phorbol ester activation (50 nM) and the adherent cells were quantitated by phosphatase assay. THP-1 cells were allowed to bind to immobilized fibrinogen (in **FIG. 8B**), or recombinant ICAM-1-Fc (in **FIG. 8C**), in the presence or absence of 200 μ M soluble peptides. All samples were assayed as triplicates and results show means \pm SD. Identical results were obtained in two other independent 25 experiments.

FIG. 8D, THP-1 cells were incubated in the presence or absence of peptides at 200 μ M concentration for 48 hours. Aliquots of conditioned medium were analyzed by gelatin zymography. Arrows show the 92 kDa proMMP-9 and 220 kDa proMMP-9 dimer.

FIG. 9A to 9C. Peptide inhibition of THP-1 cell migration. DDGW is SEQ ID NO: 174 and KKGW is SEQ ID NO: 13. THP-1 cells were preincubated with the peptide as indicated at a 200 μ M concentration for 1 h at room temperature and applied to transwells in the absence (**FIG. 9A**), or presence (**FIG. 9B**), of LLG-C4-GST coating.

Cells were allowed to migrate for 16 hours at +37°C. Cells migrated to the lower surface of the filter were stained and counted microscopically.

FIG. 9C, HT1080 fibrosarcoma cell migration was similarly assayed in the absence of LLG-C4-coating. The bars show means ± SD from triplicate wells.

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FIG. 10A to 10D. α_M -I domain binding to recombinant MMP-9 domains. HFDDDE is SEQ ID NO: 10, DFEDHD is SEQ ID NO: 21, DDGW is SEQ ID NO: 174 and KKGW is SEQ ID NO: 13.

FIG. 10A, Schematic representation of MMP-9 and its recombinant forms produced in 10 *E. coli*.

FIG. 10B, ProMMP-9, its recombinant forms or BSA were coated on microtiter wells (80 μ g/well) and soluble GST- α_M I domain was allowed to bind at the concentrations indicated. The binding was determined by anti-GST monoclonal antibody. The results are means ± SD from triplicate wells in this and other figures.

FIG. 10C, Binding of proMMP-9 to the immobilized GST- α_M I domain was studied in the presence of each peptide at the concentrations indicated. The binding was determined with the anti-MMP-9 antibody GE-213.

FIG. 10D, Binding of GST- α_M I domain to the immobilised proMMP-8, proMMP-9, ICAM-1, and fibrinogen was studied with ICAM-1, DDGW (SEQ ID NO: 174) or 20 KKGW (SEQ ID NO: 13) (50 μ M) as competitors. In control wells, GST was added instead of GST- α_M I domain. The experiment was repeated three times with similar results.

FIG. 11A to 11D. Recognition of recombinant MMP-9 domains by $\alpha_M\beta_2$ integrin-expressing cells. DDGW is SEQ ID NO: 174, KKGW is SEQ ID NO: 13, HFDDDE is SEQ ID NO: 10, and DFEDHD is SEQ ID NO: 21. The studied cells were PMNs (11A, 11B, 11C), $\alpha_M\beta_2$ L-cell transfectants (11D), non-transfectants (11D), and LAD-1 cells (11D). PMNs were in resting state or stimulated with PMA (11A, 11C) or C5a or TNF α (11B) before the binding experiment to proMMP-9 or its domains. Cells were 30 also pretreated with each peptide (50 μ M), antibody (20 μ g/ml) or the α_M I domain as indicated. Unbound cells were removed by washing and the number of adherent cells was quantitated by a phosphatase assay. The experiment was repeated three times with similar results.

FIG. 12A to 12 D. Blockage of PMN and THP-1 cell migration *in vitro* by gelatinase and β_2 integrin inhibitors.

DDGW is SEQ ID NO: 174, KKGW is SEQ ID NO: 13, HFDDDE is SEQ ID NO: 10, and DFEDHD is SEQ ID NO: 21.

- 5 PMNs (1×10^5 in 100 μ l) were applied on the LLG-C4-GST or GST coated surface (12A) or HMEC monolayer (12B) in the absence or presence of peptides (200 μ M) or antibodies (20 μ g/ml) as indicated. PMNs were stimulated with 20 nM PMA (12A), HMECs with 50 μ M C5a or 10ng/ml TNF α or left untreated (12B). THP-1 cells (5×10^4 in 100 μ l) were stimulated with 50 nM PMA and applied on the coated surfaces together
10 with each peptide (200 μ M) (12C). The cells migrated through transwell filters were stained and counted microscopically. All experiments were repeated at least twice.
(12D) Phorbol ester-activated THP-1 cells (5×10^4 in 100 μ l) were incubated for 16 h at +37°C in the presence or absence of peptides as indicated. The conditioned medium was analyzed by gelatin zymography.

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FIG. 13A to 13D. Inhibition of neutrophil migration to an inflammatory tissue.

DDGW is SEQ ID NO: 174, KKGW is SEQ ID NO: 13, HFDDDE is SEQ ID NO: 10, and DFEDHD is SEQ ID NO: 21.

- 20 FIG. 13A, Mice were injected with thioglycolate or PBS intraperitoneally. The peptides were applied intravenously at the amounts indicated. After 3 h, the intraperitoneal leukocytes were harvested and counted. The results show means \pm SD of 2–4 mice in a group. (*) indicates statistical significant difference ($p < 0.001$). The experiment was repeated at least 3 times. The infiltrated neutrophils of mice treated with thioglycolate (13B) or PBS (13C) were stained with anti-MMP-9 and anti- α_M by incubating the cells
25 with the antibodies for 3 h. Fluorescence was studied by confocal microscopy. Bars: 9.1 μ m and 4.8 μ m, respectively.

- FIG. 13D, Gelatinolytic activity of the supernatants from the peritoneal cavities of mice collected as in (13A). Lanes 1–4: samples are from thioglycolate-treated mice; lane 5: a sample from PBS-treated mouse. DGDW (SEQ ID NO: 174), HFDDDE (SEQ ID NO: 10), and DFEDHD (SEQ ID NO: 21) were injected intravenously at doses of 0.1, 0.2 and 0.2 mg per mouse. The arrows show proMMP-9 dimer, proMMP-9 and proMMP-2. The experiment was repeated three times with similar results.

The publications and other materials referred to or used herein to illuminate the

background of the invention, and in particular, to provide additional details with respect to its practice, are incorporated herein by reference. The invention will be described in more detail in the following Experimental Section.

5 EXPERIMENTAL

Antibodies and Reagents

The antibodies MEM170 and LM2/1 were against the α_M and the MEM-83 and TS2/4 antibodies against the α_L integrin subunit (19, 20). The monoclonal antibody 7E4 (21) reacted with the common β_2 -chain of the leukocyte integrins. The α_M antibody OKM10 was obtained from the American Type Culture Collection, ATCC, Rockville, MD (22). A monoclonal antibody against ICAM-5 (TL3) (23) was used as an antibody control. The monoclonal anti-MMP-9 antibody (GE-213) and anti-MMP-2 antibody (Ab-3) were obtained from Lab Vision Corporation (Fremont CA) and from OncogeneTM research products, respectively. Affinity purified rabbit anti-MMP-9 polyclonal antibodies were from the Borregaard laboratory (24). As monoclonal antibody controls, we used a mouse IgG (Silenius, Hawthorn, Australia) and anti-glycophorin A (GPA) (ATCC). Anti-trypsinogen-2 (TAT-2) antibody was a rabbit polyclonal antibody control (32). The peroxidase-conjugated anti-GST mAb was from Santa Cruz Biotechnology. The rabbit anti-mouse horseradish peroxidase-conjugated secondary antibody was from Dakopatts a/s (Copenhagen, Denmark). Inh1 (2R-2-(4-biphenylsulfonyl)amino-N-hydroxy-propionamide) was purchased from Calbiochem, La Jolla, CA. Human recombinant ICAM-1 was obtained commercially by R&D systems (Minneapolis, MN). ICAM-1-Fc fusion protein was expressed in Chinese hamster ovary cells and purified as described (14). The synthetic peptides CTT, STT, LLG-C4 and RGD-4C were obtained as previously described (14, 25). W→A CTT was ordered from Neosystem, Strasbourg, France. ProMMP-2 and proMMP-9 were obtained commercially (Roche). In zymography, the commercial proMMP-9 showed the 92kDa monomer, 200 kDa homodimer, and 120 kDa NGAL complex bands. The integrins $\alpha_1\beta_1$ and $\alpha_3\beta_1$ were purchased from Chemicon International (Temecula, CA). Human plasma fibrinogen and lovastatin were from Calbiochem.

Phage display

Phage display selections were made using a pool of random peptides CX₇₋₁₀C and X₉₋₁₀, where C is a cysteine and X is any amino acid (14, 25). Briefly, α_M I domain-GST or GST fusion protein was immobilized on microtiter wells at 20 μ g/ml concentrations and the wells were blocked with BSA. The phage library pool was first subtracted on wells 5 coated with GST and then unbound phage was transferred to α_M I domain-GST-coated wells in 50 mM Hepes/5 mM CaCl₂/1 μ M ZnCl₂/150 mM NaCl/2% BSA (pH 7.5). After three rounds of subtraction and selection, individual phage clones were tested for binding specificity and the sequences of the phage that specifically bound to the I domain were determined (14).

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Peptide biosynthesis and chemical synthesis

The phage peptides were initially prepared biosynthetically as intein fusions. The DNA sequences encoding the peptides were PCR cloned from 1 μ l aliquots of the phage-containing bacterial colonies that were stored at -20 °C. The forward primer was 5'-15 CCTTTCTGCTCTTCCAACGCCGACGGGGCT-3' (SEQ ID NO: 22) and the reverse primer 5'-ACTTTCAACCTGCAGTTACCCAGCGGCC-3' (SEQ ID NO: 23). The PCR conditions included initial denaturation at 94°C for 2 min followed by 30 cycles of 94°C 30 sec, 55°C 30 sec, and 72°C 30 sec. The PCR products were purified using QIAGEN Nucleotide removal kit. They were then digested with SapI and PstI 20 restriction enzymes and ligated to a similarly digested and phosphatase treated pTWIN vector (New England Biolabs). Correct insertions were verified by DNA sequencing. Intein fusion proteins were produced in *E. coli* strain ER2566 and affinity purified on a chitin column essentially as described (26). The peptide was cleaved on the column, eluted and finally purified by HPLC. Chemical peptide synthesis was done using Fmoc-chemistry as described and the sequences were verified by mass spectroscopy (26).

Phage binding assay

Phage (10⁸ infective particles/well) in 50 mM Hepes/5 mM CaCl₂/1 μ M ZnCl₂/0.5% BSA (pH 7.5) were added to microtiter wells coated with I domain-GST fusion or GST 30 (20 ng/well). The phages were allowed to bind in the absence or presence of a competitor peptide (15 μ M) for one hour followed by washings with PBS containing 0.05% Tween 20. The bound phage was detected using 1:3000 dilution of a peroxidase-labelled monoclonal anti-phage antibody (Amersham Biosciences) and o-phenylenediamine dihydrochloride as a substrate. The reactions were stopped by

addition of 10% H₂SO₄ and the absorbance was read at 492 nm using a microplate reader.

Pepspot

- 5 The peptides were synthesized on cellulose membranes as described (27). The membrane was blocked with 3% BSA in TBS containing 0.05% Tween 20, and incubated with 0.5-5 µg/ml α_M I domain for 2 h at room temperature. The DDGW (SEQ ID NO: 174) peptide was used as a competitor at a 50 µM concentration. Bound α_M I domain was detected using the monoclonal antibody LM2/1 (1 µg/ml) or MEM-170 (5 µg/ml) and peroxidase-conjugated rabbit anti-mouse antibody (1:5000 dilution) followed by chemiluminescence detection.
- 10

Cell culture

- The human HT1080 fibrosarcoma and THP-1 and Jurkat leukemic lines were obtained
15 from ATCC and maintained as described previously (20, 25, 28). OCI/AML-3, derived from the primary blasts of an AML patient (29) was maintained in 10% FBS/RPMI supplemented with L-glutamine, penicillin and streptomycin. Cell viability was assessed with a MTT (3-[4,5-dimethylthiazol-2-yl]-2,5-diphenyl tetrazolium bromide) assay according to the instructions of the manufacturer (Roche).

20

Purification of integrins

- $\alpha_L\beta_2$ (CD11a/CD18, LFA-1), $\alpha_M\beta_2$ (CD11b/CD18, Mac-1) and $\alpha_X\beta_2$ (CD11c/CD18) integrins were purified from human blood buffy coat cell lysates by adsorption to the anti-CD11a (TS 2/4), anti-CD11b (MEM170), or anti-CD11c (3.9) antibodies linked to
25 protein A-Sepharose CL 4B. The integrins were eluted at pH 11.5 in the presence of 2 mM MgCl₂, and 1% n-octyl glucoside as described previously (28).

Expression and purification of GST fusion proteins

- The α_L , α_M , and α_X I domains were produced as GST fusion proteins in *E. coli* strains
30 BL 21 or JM109 and purified by affinity chromatography on glutathione-coupled beads (30, 31). GST containing CTT in the C-terminus was constructed using the protocols described for LLG-C4-GST (14) and glutathione-coupled beads were employed for purification. The purity of the GST-fusion proteins was confirmed by SDS-PAGE with

Coomassie Blue staining and Western blot analysis. For pepspot analysis, GST was cleaved from the α_M I domain with thrombin.

Binding of MMPs to purified integrins

- 5 The purified I domains (GST- α_M , GST- α_L , GST- α_X), or integrins ($\alpha_M\beta_2$, $\alpha_L\beta_2$, $\alpha_X\beta_2$, $\alpha_1\beta_1$) (1 μ g/well) were immobilized in 20 mM Tris, 150 mM NaCl, 1 mM CaCl₂, 1 mM MgCl₂, and 1 mM MnCl₂, pH 7.4. The wells were washed with PBST (10 mM phosphate, 140 mM NaCl, pH 7.4, containing 0.05% Tween20) and blocked with 3% BSA in PBST. ProMMP-2, proMMP-9, or the p-aminophenyl mercuric acetate (APMA)
 10 or trypsin-activated forms (32) were incubated for 2 h at room temperature. In the inhibition experiments, CTT and Inh1 were first preincubated with the proMMPs for 30 minutes at room temperature. The wells were washed three times and incubated with anti-MMP-9 (GE-213) or anti-MMP-2 (Ab-3) antibody at a 2 μ g/ml concentration in PBST for 1 h. Bound antibodies were detected using peroxidase-conjugated rabbit anti-
 15 mouse IgG (DAKO, Glostrup, Denmark) and o-phenylenediamine dihydrochloride as a substrate.

Coprecipitation of β_2 integrin and progelatinases

- Serum-free conditioned medium containing proMMP-2 and proMMP-9 was collected
 20 from human HT-1080 fibrosarcoma cells grown in the presence of 100 nM phorbol ester 4 β -Phorbol 12,13-dibutyrate (PDBu) (Sigma-Aldrich, St. Louis, MO) overnight at +37°C. A 500 μ l volume of the supernatant was incubated with 100 ng of GST- α_M , GST- α_L , or GST- α_X I domain or $\alpha_M\beta_2$ integrin for 3 h at 25°C. GST and GST-LLG-C4 were used to determine non-specific binding. CTT, STT, LLG-C4, and ICAM-1 were
 25 used as competitors at a 200 μ g/ml concentration, and the antibodies LM2/1 and TL3 at 40 μ g/ml. After an hour incubation at +4°C, complexes of I domain and gelatinases were pelleted with Glutathione Sepharose. Integrin complexes were captured by incubating first with the OKM10 antibody for 3 h at +4°C and then with protein G Sepharose for 1 h. After centrifugation and washing, samples were analyzed by gelatin
 30 zymography on 8% SDS-polyacrylamide gels containing 0.2 % gelatin (32).

Effect of peptides on proMMP-9 release from cells

THP-1 cells (40 000/100 μ l) were incubated in serum-free RPMI medium for 48 h in the

absence or presence of 200 μ M peptide as described in the text. Aliquots of the conditioned media were analyzed by gelatin zymography.

Interaction between CTT and proMMP-9

- 5 CTT-GST and GST control (5 μ g/well) were coated overnight on 96-well microtiter plates in 50 μ l TBS followed by blocking of the wells by BSA. proMMP-9 or APMA-activated form (80 ng/well) was incubated in the absence or presence of competitors for 2 h in 50 μ M Hepes buffer containing 1 % BSA, 5 mM CaCl₂, and 1 μ M ZnCl₂ (pH 7.5). After washing, bound MMP-9 was determined with anti-MMP-9 and HRP-conjugated anti-mouse IgG as described above. To examine complexing of CTT with proMMP-9 in cell culture, THP-1 cells were activated with PDBu for 30 min and then incubated with CTT, W→A CTT, or Inh1 (each 200 μ M) at +37°C in serum-free medium. Samples were taken from the media at 0, 1, 2, 3, 4, and 5 h time points and analyzed by zymography and Western blotting with polyclonal anti-MMP-9 antibodies.
- 10 Experiments with HT-1080 cells were performed similarly except that the medium samples were collected after 6 h.
- 15

Cell surface labelling, immunoprecipitation and immunoblotting

- 20 Non-activated or PDBu-activated THP-1 cells (1×10^7) were subjected to surface labelling using periodate tritiated sodium borohydride (33). The [³H]-labelled cells were lysed with 1% (v/v) Triton X-100 in PBS, clarified by centrifugation and precleared with protein G-Sepharose. The lysate was immunoprecipitated with polyclonal anti-MMP-9, α_M (OKM-10) or β_2 (7E4) antibodies. After one-hour incubation at +4°C together with protein G-Sepharose, immunocomplexes were pelleted, washed and resolved on 8-16% SDS-PAGE gels (Bio-Rad, Hercules CA). The gels were treated with an enhancer (Amplify, Amersham Biosciences), dried and exposed. Non-labelled THP-1 cells (1×10^7) were similarly lysed and immunoprecipitated as above. The samples were resolved on 4–15% SDS-PAGE gels and transferred to nitrocellulose membranes. Immunodetection was performed with α_M (MEM170) antibody (10 μ g/ml) followed by peroxidase-conjugated anti-mouse IgG and chemiluminescence detection (Amersham Biosciences). The membranes were stripped of bound antibodies and reprobed with monoclonal α_L chain (TS2/4) or polyclonal anti-MMP-9 antibodies.
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- 30

Immunofluorescence

Immunofluorescence was performed on resting cells or the cells activated with PDBu for 30 min. A portion of the cells was treated with ICAM-1 or CTT to block β_2 integrins or gelatinases, respectively. Cells were bound to poly-L-Lysine coated cover slips, fixed with methanol for 10 min at -20°C or with 4% paraformaldehyde for 15 min at +4°C,
5 and permeabilized with 0.1% Triton X-100 in PBS at room temperature for 10 min followed by several washings. The cover slips were incubated with rabbit anti-MMP-9 polyclonal and mouse anti- α_M (OKM-10) antibodies diluted 1:500. After washing with PBS, the secondary antibodies, rhodamine (TRITC)-conjugated porcine anti-rabbit or FITC-conjugated goat anti-mouse (Fab')₂ (Dakopatts a/s, Copenhagen, Denmark) were
10 incubated at a 1:1000 dilution for 30 min at room temperature. The samples were mounted with moviol, incubated in the dark for 2 days, and examined by a confocal microscope (Leica multi band confocal imagine spectrophotometer) at a 400x magnification or a fluorescence microscope (Olympus Provis 70) at a 60x magnification.

15

Neutrophil preparations and cell lines

PMNs were isolated from peripheral blood anticoagulated in acid-citrate dextrose. Erythrocytes were sedimented by centrifugation on 2% Dextran T-500, and the leukocyte-rich supernatant was pelleted, resuspended in saline and centrifuged on a
20 Lymphoprep (Nyegaard, Oslo, Norway) at 400g for 30 minutes to separate polymorphonuclear cells from platelets and mononuclear cells (16). PMN purity was >95% with typically <2% eosinophils. Cell viability was measured using an MTT (3-[4,5-dimethylthiazol-2-yl]-2,5-diphenyl tetrazolium) bromide assay as instructed by the manufacturer (Roche).

25

Human microvascular endothelial cells (HMEC-1) (17), kindly provided by S. Mustjoki (Haartman Institute, University of Helsinki), were grown in RPMI 1640 in the presence of 10% FBS containing 2 mM glutamine, 100 IU/ml penicillin and 100 μ g/ml streptomycin. Human monocytic THP-1 cells were maintained as described (14, 25).
30 Leukocyte adhesion deficiency type-1 (LAD-1) cells, wild type and $\alpha_M\beta_2$ -transfected L929 mouse fibroblastic cells were generous gifts from Dr. Jean-Pierre Cartron (INSERM, Paris, France). These cells were maintained as described previously (18) and the $\alpha_M\beta_2$ expression was examined by fluorescence-activated cell sorting (FACS, Becton Dickinson, San Jose, CA).

Cell adhesion and migration

Fibrinogen and ICAM-1-Fc were coated at 40 µg/ml in TBS at +4°C. Peptides (2 µg/well) were coated in TBS containing 0.25% glutaraldehyde at +37°C. The wells
5 were blocked with 1% BSA in PBS. THP-1 cells (50 000/well) with or without PDBu activation were added in 0.1% BSA-RPMI medium in the presence or absence of 200 µM peptides or monoclonal antibodies at 50 µg/ml. After 30-35 minutes the wells were washed with PBS to remove non-adherent cells and the adhesive cells were quantitated by a phosphatase assay. The cell migration assay was conducted using transwell
10 migration chambers (8 µm pore size, Costar) in serum-containing medium as described (14). Briefly, the membranes were coated on the upper and lower surface with 40 µg/ml GST, LLG-C4-GST, or left uncoated. The wells were blocked with 10% serum-containing medium for 2 h. THP-1 cells (50 000/100 µl) or HT1080 (20 000/100 µl)
15 were preincubated with the peptides for 1 h before transfer to the upper chamber. The lower chamber contained 500 µl of the medium without the peptides. The cells were allowed to migrate to the lower surface of the membrane for 16 h and then stained with crystal violet and counted.

MMP-9 proteins (200 nM in PBS) were coated at +4°C for 16 h and the microtiter wells
20 were blocked with 3 % BSA in PBS for 1 h at room temperature. The α_Mβ₂-integrin L-cell transfectants and PMNs (1x10⁵ cells/well) were suspended in RPMI medium supplemented with 2mM MgCl₂ and 0.1% BSA and activated with PMA (20nM) for 20 min, or with C5a (50nM) or TNF-α (10nM) for 4h at +37°C. The L926 wild type and LAD-1 cells were used as controls. The cells were treated with the indicated antibody
25 (20 µg/ml) or peptide (50 µM) at +37°C for 30 min, washed twice with serum-free medium and incubated in the microtiter wells at +37°C for 30 min. The wells were washed with PBS, and the number of adherent cells was quantitated by a phosphatase assay (14).

30 Cell migration was conducted using Costar 24-transwell migration chambers with a 3 µm pore size for PMNs and 8 µm for THP-1 cells. To study β₂ integrin-directed migration, the chamber membrane was coated on both sides with LLG-C4-GST integrin ligand (40µg/ml) or GST as a control and blocked with 10% serum-containing medium as described above. To study transendothelial migration, confluent HMECs (4x10⁵

cells/well) were grown on the upper side of the gelatin-coated membrane for 5 days. Culture medium was changed after 3 days. After washing the HMEC layers twice with PBS, chemotactic activation was carried out by adding C5a (50nM), TNF- α (10ng/ml), or medium alone to the lower compartment at +37°C for 4 h. Cultures were then washed 5 again twice to remove all agents. PMNs or THP-1 cells were preincubated with the peptide inhibitor or antibody studied for 1 h before transfer to the upper compartment (1x10⁵ cells in 100 μ l RPMI/0.1 % BSA or the complete 10 % FCS-containing medium). PMNs were allowed to migrate for 2 h through the LLG-C4-GST coated membrane and for 30 min through the HMEC monolayer. THP-1 cells were allowed to migrate for 16 10 h. The non-migrated cells were removed from the upper surface by a cotton swab and the cells that had traversed the filters were stained with crystal violet and counted.

Mouse inflammation model

Balb/c mice at the age of 31-32 weeks were injected intraperitoneally with 3% (w/v) 15 thioglycolate in sterile saline (36). Peptides (5–500 μ g in 100 μ l) were introduced intravenously through the tail vein. Animals were euthanized after 3 h and the peritoneal cells were harvested by injecting 10 ml of sterile PBS through the peritoneal wall. Red blood cells present in the lavage fluid were removed by hypotonic lysis. Cells were centrifuged and resuspended in 1 ml of sterile 0.25% BSA/Krebs-Ringer. The 20 supernatants were also collected and analysed by gelatin zymography. The number of neutrophils was determined following staining with 0.1% crystal violet and using a light microscope equipped with a x 100 objective. For immunofluorescence staining, cells were allowed to bind to poly-L-lysine coated cover slips, fixed with 2.5% paraformaldehyde in PBS at +4°C for 30 min followed by several washings. The Fc 25 receptors were blocked in the presence of 20% of rabbit serum and 3% BSA in PBS. The cells were then incubated with anti-MMP-9 polyclonal and α_M monoclonal (MCA74) antibodies for 30 min. After washing with PBS, the secondary antibodies, rhodamine (TRITC)-conjugated anti-rabbit or FITC-conjugated anti-rat (Fab')₂ were 30 incubated for another 30 min. The samples were examined with a confocal microscope. The animal studies were approved by an ethical committee of Helsinki University.

RESULTS

Identification of the α_M I domain-binding peptide motif D/E-/D/E-G/L-W (*SEQ ID*

NO: 2)

Using phage peptide display libraries, we selected peptides that interact with the α_M I domain. GST-binding phage were first eliminated on GST-coated wells and the unbound phage preparations were incubated on α_M I domain GST fusion protein-coated wells. The α_M I domain-binding phage were enriched by three rounds of panning and the peptide sequences were determined. With the exception of one linear peptide, the peptides were derived from the cyclic CX₇C and CX₈C libraries. The I domain-binding sequences showed only one conserved motif, a somewhat unexpected finding in terms of the known ligand binding promiscuity of the I domain. The bound peptides contained two consecutive negatively charged amino acids, i.e. glutamatic and/or aspartic acids, followed by glycine and tryptophan residues (Fig. 1A). The consensus D/E-/D/E-G/L-W (SEQ ID NO: 2) determined by this approach was clearly different from LLG-C4 and other β_2 integrin-binding peptides reported so far.

We first prepared the phage display peptides as intein fusion proteins, from which the peptides were cleaved. This allowed us to rapidly test the peptide solubility and the binding specificity before large-scale chemical peptide synthesis. The peptides were cloned using oligonucleotide primers that amplify the peptide library insert from the phage vector. Consequently, all the peptides prepared contain the vector-derived sequences ADGA (SEQ ID NO: 24) and GAAG (SEQ ID NO: 25) in the NH₂- and COOH- termini, respectively. Phage binding experiments using soluble peptides as competitors indicated that the peptides bearing the two adjacent negative charges bound to a common site (not shown). We chose the peptide ADGA-CILWMDDGWC-GAAG (SEQ ID NO: 12) (DDGW)(SEQ ID NO: 174) for further experiments as this peptide showed strong binding and was highly soluble in aqueous buffers (soluble in 50 mM NaOH at >10 mM concentrations). The peptide was also prepared by chemical synthesis. The phage bearing the DDGW (SEQ ID NO: 174) sequence avidly bound to the α_M I domain and this was readily inhibited by low concentrations of the DDGW (SEQ ID NO: 174) peptide, but only marginally affected by the LLG-C4 peptide, indicating different binding sites for DDGW (SEQ ID NO: 174) and LLG-C4 (Fig. 1B). Control phage bearing other peptide sequences did not bind. The DDGW (SEQ ID NO: 174) bearing phage also showed also specific binding to the α_L I domain that was inhibitable by DDGW (SEQ ID NO: 174) but the interaction was weaker than with the

α_M I domain (Fig. 1C and data not shown). No binding was observed with the α_X I domain or GST used as a control (Fig. 1C).

Characterization of DELW (SEQ ID NO: 175) sequence on the catalytic domain of gelatinases that mediates interaction with the β_2 integrin I domains

We searched protein databases for matches to the novel D/E-/D/E-G/L-W (SEQ ID NO: 2) motif. One of the phage library-derived peptides, CPEELWWLC (SEQ ID NO: 19), was highly similar to the DELW(S/T)LG (SEQ ID NO: 8) sequence present on the catalytic domain of MMP-2 and MMP-9 gelatinases (Fig. 1A). DELW (SEQ ID NO: 175) like sequences with double negative charges are also present in other secreted MMPs but not in the membrane-type MMPs such as MMP-14.

No MMP has been reported to bind to the leukocyte β_2 integrins. We therefore set out to study whether MMP-9 in particular could be a ligand of the β_2 integrins as MMP-9 gelatinase is the major leukocyte MMP and is induced during β_2 integrin activation. As a first step, we synthesized the whole proMMP-9 sequence as overlapping 20-mer peptides on a pepspot membrane. Binding assays with the α_M I domain revealed a single active peptide that located to the MMP-9 catalytic domain (Fig. 1D). No binding was observed, when the I domain was omitted and the membrane was probed with antibodies only. The sequence of the I domain-binding peptide was QGDAHFDDDELWSLGKGVVV (SEQ ID NO: 26) and it contained the binding motif identified by phage display (Fig. 1D).

The active MMP-9 peptide contained four consecutive amino acids with negative charges, DDDE (SEQ ID NO: 27). To study the importance of these residues, the aspartic and glutamic acid residues that were closest to the tryptophan were replaced by alanines. At the same time the peptide length was shortened to 15-mer. The alanine mutagenesis significantly abrogated I domain binding on the pepspot filter; the OD value dropped from 2010 to 476 (Table I). To study whether the negatively charged peptide from other MMPs is also active, we synthesized the corresponding 15-mers and the double alanine mutations. Sequences from MMP-1, 2, 3, 7, 8, 9 and 13, but not the membrane-anchored MMP-14 (MT1-MMP), could bind the α_M I domain. Alanine mutations always decreased the binding.

- We did similar pepspot analysis for some of the known I domain ligands, which contain D/E-/D/E-G/L-W (SEQ ID NO: 2) like sequences. Peptides derived from myeloperoxidase, catalase, thrombospondin-1 and complement protein iC3b strongly bound the I domain in this assay and the double alanine mutation caused a loss of binding (see Table I). Of the three iC3b peptide permutations tested, ARSNLDEDIIAEENI (SEQ ID NO: 5) was the active one. The acidic residues were followed by a hydrophobic isoleucine cluster in this peptide. The soluble DDGW (SEQ ID NO: 174) peptide efficiently inhibited the binding of this peptide to the I domain.
- 10 Weaker I domain binding was observed with one complement factor H-derived peptide and one fibronectin-derived peptide. Peptides derived from ICAMs-1, 2 and 3, neutrophil inhibitory factor, Cyr61, fibrinogen, GP1b, factor X, or E-selectin lacked activity.
- 15 Alanine scanning mutagenesis of the DDGW (SEQ ID NO: 174) peptide with the pepspot system similarly indicated the importance of the glutamic acid residues for I domain binding (Fig. 1E). Alanine mutations of the glycine or either one of the tryptophan residues also inactivated the peptide. Mutations of the isoleucine, leucine or methionine residues were tolerated. Deletion of the ADGA (SEQ ID NO: 24) sequence from the N terminus had no effect on I domain binding, but removal of the C terminal GAAG (SEQ ID NO: 25) sequence abolished the binding. As the peptides were immobilized via the C terminus on the filter, a sufficient linker sequence such as GAAG (SEQ ID NO: 25) seemed important. We also tested a series of truncated cyclic peptides to identify the shortest active sequence. This analysis showed that ADGA-CEDGWC-
- 20 GAAG (SEQ ID NO: 28), but not ADGA-CDDGWC-GAAG (SEQ ID NO: 29) was the minimal peptide that supported α_M I domain binding. The longer side chain of glutamate compared to aspartate is probably required to bring the negatively charged carboxyl group in the correct position for I domain binding.
- 25
- 30 **Progelatinases bind to purified $\alpha_M\beta_2$ and $\alpha_L\beta_2$ integrins and their I domains**
- We next used a microtiter well-based sandwich assay to study gelatinase binding to purified integrins. Progelatinases bound in a concentration-dependent manner to coated $\alpha_M\beta_2$ integrin (Fig. 2A). Curiously, MMP-2 and MMP-9 lost the integrin binding ability after activation by trypsin or APMA. The binding of proMMP-2 and proMMP-9 was

observed with both $\alpha_M\beta_2$ and $\alpha_L\beta_2$ integrins and their corresponding I domains (Fig. 2B and 2C). No binding was detected on the α_X I domain or the $\alpha_1\beta_1$ and $\alpha_3\beta_1$ integrins.

The DDGW (SEQ ID NO: 174) peptide was an efficient inhibitor and it inhibited proMMP-9 binding to the α_M I domain with an IC₅₀ of 20 μ M (Fig. 3A and 3B). To demonstrate that the negative charges of aspartic acids are essential for the peptide activity, the peptide ADGACILWMKKKGWCGAAG (SEQ ID NO: 14) (KKGW) (SEQ ID NO: 13) containing lysines in place of aspartic acids was prepared. As expected, the KKWG (SEQ ID NO: 13) peptide was inactive and did not compete with proMMP-9 binding. We were also interested in testing lovastatin, as its binding site in the α_L I domain is known (34, 35). Lovastatin was not able to compete with proMMP-9 even at a high concentration.

ProMMP-9 bound like a true integrin ligand, as the cation chelator EDTA (5 mM) nearly completely prevented the binding (Fig. 3C and 3D). For background measurement in the sandwich assay, we used antibodies alone (control), or coating with ICAM-1 or wild type GST. The gelatinase-binding peptide CTT (200 μ M) inhibited proMMP-9-integrin interaction with the same efficiency as EDTA did. The control peptides STTHWGFTLS (SEQ ID NO: 16) (STT) and CTTHAGFTLC (SEQ ID NO: 17) (W→A CTT), which lack gelatinase inhibitory activity (26), were without effect. A non-peptide chemical MMP inhibitor (Inh1) also prevented proMMP-9 binding. As EDTA inhibits both the gelatinase and the integrin, we used integrin blocking antibodies and ligand peptides to demonstrate the specific binding activity of β_2 integrin. The known ligand-binding blocking antibodies MEM 170, MEM 83, and LM2/1 inhibited proMMP-9 binding. A control antibody TL3 had no effect. The I domain binding peptide LLG-C4 showed a partial inhibitory effect. RGD-4C, a ligand of α_V integrins, served as control peptide and had no effect on proMMP-9 binding. The purity of the integrins was typically more than 90% and that of I domains 95%, making it unlikely that progelatinases would bind to impurities in the preparations.

30

Progelatinase-integrin complexes were also obtained by co-precipitation experiments using HT1080 conditioned medium as a source of proMMP-9 and proMMP-2, which were analyzed by zymography. The progelatinases co-precipitated with $\alpha_M\beta_2$ integrin

- or α_M I domain GST fusion protein when these were used as a bait. The integrin added to the medium was immunoprecipitated with the α_M antibody OKM10 (Fig. 4A). The α_M I domain GST protein was pulled down with glutathione-beads (Fig. 4B). CTT but not STT had an inhibitory effect. Inhibition of the I domain by LM2/1, ICAM-1 or LLG-C4 also affected the pull-down of progelatinases. GST control did not coprecipitate the gelatinases. No active forms of gelatinases were found to coprecipitate with $\alpha_M\beta_2$ or the I domain, when APMA-treated HT-1080 medium or APMA-activated MMP-9 was used (not shown).
- As the gelatinase inhibitors CTT and Inh1 prevented the binding of proMMP-9 to the integrin, it can be anticipated that CTT and Inh1 avidly bind to proMMP-9. To gain more insight into this, we examined binding of proMMP-9 to immobilized CTT peptide. ProMMP-9 specifically bound to the CTT-GST fusion protein (Fig. 5A) but not to LLG-C4-GST. CTT and Inh1 at 100 μ M concentrations effectively competed in binding but W→A CTT did not. The proMMP-9 preparation did not contain detectable amounts of active MMP-9 on zymography analysis, and after proMMP-9 activation with APMA, the CTT-GST binding increased. CTT and Inh1 could also bind to proMMP-9 secreted into the medium of PDBu-activated THP-1 leukemic cells (Fig. 5B) or HT1080 fibrosarcoma cells (not shown). A time-dependent reduction in the gelatinolytic activity of proMMP-9 was observed with CTT (panel 1) and Inh1 (panel 3), but not with the W→A CTT peptide (panel 4). Western blot analysis indicated that CTT does not decrease the secretion of proMMP-9 by the cells (panel 2). Furthermore, the CTT complex was reversible and disappeared after repeated freezing and thawing of the samples.

25

Demonstration of a cell-surface complex between progelatinases and β_2 integrins

To study whether the progelatinases occur in a complex with the β_2 integrins on the leukocyte surface, we performed immunoprecipitation and co-localization studies. First, we examined THP-1 monocytic leukemia cells in the resting state and after induction by PDBu, which mimics leukocyte activation *in vivo*. THP-1 cell-stimulation with PDBu led to upregulation of MMP-9 (data not shown). The cell surface glycoproteins of THP-1 cells were labelled with tritium [3 H] followed by immunoprecipitation with β_2 integrin and MMP-9 antibodies. In the PDBu-activated cells, the α_M chain antibody OKM10 and

β₂ chain antibody 7E4 immunoprecipitated two [³H]-labelled proteins corresponding to the integrin α_M chain (165 kDa) and β₂ chain (95 kDa) (Fig. 6A, lanes 9-10). Importantly, polyclonal MMP-9 antibodies immunoprecipitated the same two integrin chains (lane 7). In non-activated cells, essentially no co-precipitation of α_M and β₂ were observed with MMP-9 antibodies, although the α_M and β₂ chains were present. The co-precipitation of the integrin chains by MMP-9 antibodies was prevented by the CTT peptide (lane 8). The control antibody (TL3) did not precipitate any proteins.

With the [³H]-labelled cells, we did not observe any band corresponding to proMMP-9, perhaps because the carbohydrates of proMMP-9 are poorly labelled. We therefore analyzed the PDBu-activated THP-1 cells by Western blotting (Fig. 6B). ProMMP-9 was readily immunoprecipitated with antibodies against MMP-9, α_M or α_L, but not by the control antibody. MMP-9 antibodies in turn were able to immunoprecipitate the α_M but not the α_L chain. MMP-2 antibodies similarly co-precipitated α_M but not α_L. When the cell lysate was precleared with the α_M antibody OKM-10, the amount of immunoprecipitated α_M and proMMP-9 clearly decreased (lane 6). Preclearing with the α_L antibody TS2/4 did not significantly remove α_M or proMMP-9, but abolished the α_L precipitation (lane 7).

As THP-1 cells do not express high amounts of the α_L chain (20), we examined the Jurkat T cell line, which expresses more α_L than α_M (28). We observed a significant immunoprecipitation of α_L by MMP-9 and MMP-2 antibodies after PDBu activation (Fig. 6C). Furthermore, the α_L antibody co-precipitated more proMMP-9 in comparison to the α_M antibody. No proMMP-9 co-precipitated with MMP-2 antibodies in Jurkat or THP-1 cells.

ProMMP-9 and α_Mβ₂ were found to co-localize on the cell surface following PDBu-activation of THP-1 cells as studied by fluorescence and confocal microscopy (Figs. 7A and 7B, respectively). Using a higher magnification, colocalization was primarily seen in cell surface clusters (Fig. 7B), and to a lesser extend on areas where cells contacted each other (not shown). We believe that the MMP-9 co-localizing with α_Mβ₂ is the proMMP-9, as the activated MMP-9 did not bind to α_Mβ₂. Without PDBu activation,

there was hardly any co-localization of proMMP-9 and $\alpha_M\beta_2$. The secondary antibodies did not stain the cells when the primary antibodies were omitted (data not shown).

When the cells were preincubated with the CTT peptide or recombinant soluble ICAM-1 to block proMMP-9 or $\alpha_M\beta_2$, the cell surface clusters did not form and the proMMP-

- 5 9- $\alpha_M\beta_2$ colocalization was not observed (not shown). ProMMP-9- $\alpha_M\beta_2$ colocalization was also observed on Jurkat cells following phorbol ester stimulation (not shown).

Blocking the progelatinase/ β_2 integrin complex with DDGW (SEQ ID NO: 174) releases cell-bound proMMP-9 and inhibits cell migration but not adhesion

- 10 As the DDGW peptide is an integrin ligand, one of the questions was whether it can support adhesion of leukocytes. We studied adhesion of human myelomonocytic THP-1 cells on immobilized glutaraldehyde-polymerized peptide. Phorbol-ester activated cells efficiently bound to the DDGW peptide, whereas there was no binding in the absence of cell activation (Fig. 8A). As a positive control, the recombinant intein-produced ADGA-
- 15 CPCFLLGCC-GAAG (SEQ ID NO: 30) peptide supported adhesion, but unlike the DDGW peptide, it also supported adhesion in the absence of integrin activation. The acute myeloid leukemic cell line OCI/AML-3 also avidly adhered to DDGW (SEQ ID NO: 174), whereas human fibrosarcoma HT1080 cells which lack β_2 integrins did not (not shown). As THP-1 cells were able to adhere on DDGW (SEQ ID NO: 174), we
- 20 next studied the effect of the peptide on β_2 integrin dependent adhesion to fibrinogen and ICAM-1. Interestingly, DDGW did not block cell adhesion to fibrinogen, whereas the LLG-C4 peptide blocked the adhesion as previously reported (Fig. 8B). Similarly, DDGW (SEQ ID NO: 174) did not block the binding of recombinant α_M I domain to immobilized fibrinogen (not shown). DDGW did not either block cell adhesion on
- 25 ICAM-1-Fc fusion protein. As a control, the blocking antibody 7E4 against β_2 integrins prevented the ICAM-1 binding, indicating that the THP-1 cells bound in a β_2 integrin dependent manner (Fig. 8C). We also found no blocking effect of DDGW (SEQ ID NO: 174) on THP-1 adhesion to LLG-C4-GST fusion protein (not shown).
- 30 The second question raised by these studies was whether the DDGW (SEQ ID NO: 174) peptide can release cell-bound proMMP-9. When THP-1 cells were cultured for 48 h in the presence of DDGW (SEQ ID NO: 174), an increase of proMMP-9 level was observed in the conditioned medium as studied by gelatin zymography (Fig. 8D). The

peptide increased both monomeric and dimeric proMMP-9 in the culture medium. In contrast, CTT slightly decreased or inhibited active proMMP-9. KKGW (SEQ ID NO: 13) and W→A CTT had no effect.

5 We also studied the role of the progelatinase/ β_2 integrin complex in leukocyte migration using a transwell assay in which leukocyte migration can be adjusted by the choice of coated matrix or ligand protein. We tested that the CTT, LLG and DDGW (SEQ ID NO: 174) peptides are not toxic to the THP-1 cells in a 48 h time frame at >200 μ M concentrations using an MTT assay. Using transwells coated with 10% serum in cell
10 culture medium, we first studied the effect of peptides on the basal migration of THP-1 cells in the absence of any stimulus by phorbol ester or an adhesive matrix. Under such conditions, CTT, LLG-C4 or the gelatinase inhibitor Inh1 at a 200 μ M concentration had no effect on THP-1 migration indicating no active involvement of gelatinases or β_2 integrins (Fig. 9A). We have previously shown that when the transwells are coated with
15 LLG-C4-GST fusion protein, THP-1 cells adhere and migrate in a β_2 integrin dependent manner (14). Thus, transwells were coated with LLG-C4-GST fusion protein or GST alone. Both the DDGW (SEQ ID NO: 174) and CTT peptide, but not KKGW (SEQ ID NO: 13), inhibited the migration of THP-1 cells on the LLG-C4-GST substratum (Fig. 9B). The soluble LLG-C4 peptide also blocked the migration. In the presence of GST
20 coating, cell migration was negligible. To verify that the effect of DDGW peptide was β_2 integrin dependent, HT1080 fibrosarcoma cells lacking these integrins were allowed to migrate in the presence of CTT, DDGW (SEQ ID NO: 174), KKGW (SEQ ID NO: 13) or LLG-C4. Of these peptides, only CTT was capable of inhibiting cell migration (Fig. 9C).

25

Peptide inhibitors of the proMMP-9/ $\alpha_M\beta_2$ complex prevent neutrophil migration

As shown above, pepspot analysis located the integrin interactive site of proMMP-9 to a 20-amino acid long sequence present in the catalytic domain, QGDAHFDDDE-LWSLGKGVVV (SEQ ID NO: 26). Further screening by the pepspot system has
30 indicated that sufficient integrin binding activity is achieved by truncating this sequence to a hexapeptide, HFDDDE (SEQ ID NO: 10) (data not shown). To confirm that such a short sequence is the bioactive site of proMMP-9, we first prepared bacterially expressed recombinant domains of MMP-9 (Fig. 10A). Δ MMP-9 is composed of the

prodomain (Pro) and the catalytic domain but lacks the hemopexin domain. The fibronectin type II repeats (FnII) were also produced as a separate recombinant protein as this is an important substrate-binding region. The procatalytic domain construct Δ MMP-9 bound the α_M I domain nearly as efficiently as the wild type proMMP-9 (Fig. 10B). FnII protein almost lacked activity. The HFDDDE (SEQ ID NO: 10) peptide identified by the solid-phase pepspot analysis was highly active when made by peptide synthesis and inhibited proMMP-9 binding to the α_M I domain with an IC₅₀ of 20 μ M (Fig. 10C). The bound proMMP-9 was determined with the GE-213 antibody, which recognizes an epitope of the FnII domain (data not shown). A scrambled peptide DFEDHD (SEQ ID NO: 21) with the same set of negatively charged amino acids was inactive. HFDDDE (SEQ ID NO: 10) was equally potent as DDGW (SEQ ID NO: 174), the α_M I domain-binding peptide discovered by phage display and described above. KKGW (SEQ ID NO: 13), the control peptide for DDGW (SEQ ID NO: 174), was without effect. As the HFDDDE (SEQ ID NO: 10) sequence is highly conserved in the members of the MMP family, we also examined the α_M I domain binding to human neutrophil collagenase, MMP-8. I domain showed a similar DDGW(SEQ ID NO: 174) inhibitable binding to proMMP-8 as to proMMP-9 (Fig. 10D). ICAM-1 and fibrinogen did not compete with either proMMP, implying different binding sites for the matrix proteins and proMMPs in the I domain.

After integrin activation, PMNs exhibited an ability to adhere on proMMP-9. PMA-stimulated PMNs bound to microtiter well-coated Δ MMP-9 nearly as strongly as to proMMP-9 (Fig. 11A). Stimulation of PMNs with C5a or TNF- α gave similar results PMN adherence increasing by 3-fold (Fig. 11B). The FnII domain did not support PMN adhesion. PMN adherence was inhibited by HFDDDE (SEQ ID NO: 10) (50 μ M), DDGW (SEQ ID NO: 174) (50 μ M), the soluble α_M I domain and the MEM170 antibody (Fig. 11C), indicating β_2 integrin-directed binding. The control peptides (DFEDHD (SEQ ID NO: 21), KKGW (SEQ ID NO: 13)) and an irrelevant monoclonal antibody (anti-GPA) had no effect. The CTT peptide, but not the W→A CTT control peptide lacking MMP inhibitory activity, binds to the MMP-9 catalytic domain (unpublished results) and also inhibited the PMN adherence. MMP-9 antibodies inhibited partially.

We also examined $\alpha_M\beta_2$ -transfected L cells. The $\alpha_M\beta_2$ L-cell transfectants bound to

proMMP-9 and Δ MMP-9 similarly as PMNs did and the I domain ligands and MMP-9 inhibitors attenuated the binding (Fig. 11D). The transfected cells also showed a weak adherence to FnII domain, but the studied peptides and antibodies did not inhibit this binding. Wild type L cells or LAD-1 cells showed no binding to proMMP-9 or its domains.

The *in vitro* migration of PMNs was studied on transwell filter assays. Coating with the artificial β_2 integrin ligand LLG-C4-GST renders cell migration dependent on the β_2 integrins (14). The migration of PMA-activated PMNs was 5-fold in the LLG-C4-GST substratum in comparison to GST substratum (Fig. 12A). HFDDDE (SEQ ID NO: 10) (200 μ M) inhibited the migration of PMA-stimulated cells but not the basal migration of non-activated cells. DDGW (SEQ ID NO: 174), CTT, MEM170 (20 μ g/ml) and polyclonal anti-MMP-9 (20 μ g/ml) worked similarly, affecting the migration of the PMA-activated cells only. Control peptides and an antibody control (anti-TAT-2) had no effect. Similar results were obtained in a transendothelial migration assay (Fig. 12B). Chemotaxis with C5a or TNF- α increased PMN transmigration by 5-10 fold and inhibition was obtained by DDGW (SEQ ID NO: 174), HFDDDE (SEQ ID NO: 10), and CTT but not with the control peptides. Similarly, α_M and MMP-9 antibodies inhibited but an antibody control (anti-GPA) did not. We also examined the effects of peptides on THP-1 leukemia cell migration through the LLG-C4-GST coated transwell filters. The results were the same as for PMNs. HFDDDE (SEQ ID NO: 10), DDGW (SEQ ID NO: 174), and CTT inhibited THP-1 migration and the control peptides did not (Fig. 12C).

We show above that the DDGW (SEQ ID NO: 174) peptide can release proMMP-9 from THP-1 cells. We found that the HFDDDE (SEQ ID NO: 10) peptide also released proMMP-9 but was less effective than DDGW (SEQ ID NO: 174) (Fig. 12D). The scrambled peptide did not induce the release of proMMP-9. Under the 16 h incubation time, the peptides had no effect on the secretion of proMMP-2.

To study neutrophil migration *in vivo*, we used a mouse model of thioglycolate-induced peritonitis. The cells that infiltrated into the peritoneal cavity within 3 h after thioglycolate irritant were judged to be predominantly PMNs by crystal violet staining. The DDGW (SEQ ID NO: 174) and HFDDDE (SEQ ID NO: 10) peptides had potent *in*

vivo activities in this inflammation model (Fig. 13A). An intravenous tail injection of DDGW (SEQ ID NO: 174) or HFDDDE (SEQ ID NO: 10) inhibited the intraperitoneal accumulation of PMNs. The KKGW (SEQ ID NO: 13) and DFEDHD (SEQ ID NO: 21) peptides used as controls had no effect. The effects of DDGW (SEQ ID NO: 174) and HFDDDE (SEQ ID NO: 10) were concentration-dependent and up to 90 % inhibition was obtained by doses of 50 μ g and 500 μ g per mouse, respectively. DDGW (SEQ ID NO: 174) was active even at 5 μ g given per mouse corresponding to an effective dose of 0.1 mg/kg mouse tissue. Approximately 20-fold more PMNs were present intraperitoneally after thioglycolate-stimulus in comparison to the PBS control. The collected inflammatory PMNs stained positively for the proMMP-9/ $\alpha_M\beta_2$ complex by double immunofluorescence (Fig. 13B). The cells collected after PBS injection lacked the complex; they expressed the integrin but had no cell-surface MMP-9 (Fig. 13C). Zymography analysis of the supernatants from the collected intraperitoneal fluid showed that thioglycolate induced elevated levels of gelatinases in comparison to PBS (Fig. 13D). DDGW (SEQ ID NO: 174) and HFDDDE (SEQ ID NO: 10), but not the scrambled peptide, prevented the increase in gelatinase levels in accordance with the inhibition of cell migration.

REFERENCES

1. Gahmberg, C.G. (1997) *Curr. Opin. Cell Biol.* **9**, 643-650.
2. Colombatti, A., and Bonaldo, P. (1991) *Blood* **77**, 2305-2315.
3. Yakubenko, V. P., Lishko, V. K., Lam, S. C., and Ugarova, T. P. (2002) *J. Biol. Chem.* **107**, 1
4. Davis, G.E. (1992) *Exp. Cell Res.* **200**, 242-252.
5. Johansson, M. W., Patarroyo, M., Öberg, F., Siegbahn, A., and Nilsson, K. (1997) *J Cell Sci.* **110**, 1133-1139.
6. Cai, T. Q., and Wright, S. D. (1996) *J. Exp. Med.* **184**, 1213-1223.
7. Pluskota, E., Soloviev, D. A., and Plow, E. F. (2003) *Blood* **101**, 1582-1590.
8. Staunton, D. E., Dustin, M. L., Erickson, H. P., and Springer, T. A. (1990) *Cell* **61**, 243-254.
9. Berendt, A. R., McDowell, A., Craig, A. G., Bates, P. A., Sternberg, M. J., Marsh, K., Newbold, C. I., and Hogg, N. (1992) *Cell* **68**, 71-81.
10. Shimaoka, M., Xiao, T., Liu, J. H., Yang, Y., Dong, Y., Jun, C. D., McCormack, A., Zhang, R., Joachimiak, A., Takagi, J., Wang, J. H., and Springer, T. A. (2003) *Cell* **112**, 99-111.
11. Li, R., Xie, J., Kantor, C., Koistinen, V., Altieri, D. C., Nortamo, P., and Gahmberg, C. G. (1995) *J. Cell Biol.* **129**, 1143-1153.
12. Ugarova, T. P., Solov'ev, D. A., Zhang, L., Loukinov, D. I., Yee, V. C., Medved, L. V., and Plow, E. F. (1998) *J. Biol. Chem.* **273**, 22519-22527.
13. Schober, J.M., Lau, L.F., Ugarova, T.P., Lam, S.C. (2003) *J. Biol. Chem.* In press May 6, PMID:12736251.
14. Koivunen, E., Ranta, T. M., Annila, A., Taube, S., Uppala, A., Jokinen, M., van Willigen, G., Iharus, E., and Gahmberg, C. G. (2001) *J. Cell Biol.* **153**, 905-916.
15. Pasqualini, R., Koivunen, E., and Ruoslahti, E. (1995) *J. Cell Biol.* **130**, 1189-1196.
16. Boyum, A. (1968). *Scand. J. Clin. Lab Invest.* **97**, 77.
17. Mustjoki, S., R. Alitalo, E. Elonen, O. Carpen, C. G. Gahmberg, and A. Vaheri. (2001). *Br. J. Haematol.* **113**, 989.
18. Hermand, P., M. Huet, I. Callebaut, P. Gane, E. Iharus, C. G. Gahmberg, J. P. Cartron, and P. Bailly. (2000). *J. Biol. Chem.* **275**, 26002.
19. Li, R., Nortamo, P., Valmu, L., Tolvanen, M., Huuskonen, J., Kantor, C., and Gahmberg, C.G. (1993) *J. Biol. Chem.* **268**, 17513-17518.
20. Li, R., Xie, J., Kantor, C., Koistinen, V., Altieri, D. C., Nortamo, P., and Gahmberg, C. G. (1995) *J. Cell Biol.* **129**, 1143-1153.
21. Nortamo, P., Patarroyo, M., Kantor, C., Suopanki, J., and Gahmberg, C.G. (1988) *Scand. J. Immunol.* **28**, 537-546.
22. Wright, S.D., Rao, P.E., Van Voorhis, W.C., Craigmyle, L.S., Iida, K., Talle, M.A., Westberg, E.F., Goldstein, G., and Silverstein, S.C. (1983) *Proc. Natl. Acad. Sci. USA* **80**, 5699-5703.
23. Tian, L., Kilgannon, P., Yoshihara, Y., Mori, K., Gallatin, M., Carpen, O., and Gahmberg, C.G. (2000) *Eur. J. Immunol.* **30**, 810-818.
24. Kjeldsen, L., Johnsen, A. H., Sengelov, H., and Borregaard, N. (1993) *J. Biol. Chem.* **268**, 10425-10432.
25. Koivunen, E., Arap, W., Valtanen, H., Rainisalo, A., Medina, O. P., Heikkila, P., Kantor, C., Gahmberg, C. G., Salo, T., Konttinen, Y. T., Sorsa, T., Ruoslahti, E., and Pasqualini, R. (1999) *Nat. Biotechnol.* **17**, 768-774.
26. Björklund, M., Valtanen, H., Savilahti, H., and Koivunen, E. (2003) *Comb.*

- Chem. High Throughput Screen.* **6**, 29-35.
27. Heiskanen, T., Lundkvist, A., Soliymani, R., Koivunen, E., Vaheri, A., and Lankinen, H. (1999) *Virology* **262**, 321-332.
28. Tian, L., Yoshihara, Y., Mizuno, T., Mori, K., and Gahmberg, C. G. (1997) *J. Immunol.* **158**, 928-936.
29. Wang, C., Curtis, J. E., Minden, M. D., and McCulloch, E. A. (1989) *Leukemia* **3**, 264-269.
30. Griggs, D.W., Schmidt, C.M., and Carron, C.P. (1998) *J. Biol. Chem.* **273**, 22113-22119.
31. Michishita, M., Videm, V., and Arnaout, M.A. (1993) *Cell* **72**, 857-867.
32. Sorsa, T., Salo, T., Koivunen, E., Tyynela, J., Konttinen, Y. T., Bergmann, U., Tuuttila, A., Niemi, E., Teronen, O., Heikkila, P., Tschesche, H., Leinonen, J., Osman, S., and Stenman, U.H. (1997) *J. Biol. Chem.* **272**, 21067-21074.
33. Gahmberg, C. G., and Andersson, L. C. (1977) *J. Biol. Chem.* **252**, 5888-5894.
34. Kallen, J., Welzenbach, K., Ramage, P., Geyl, D., Kriwachi, R., Legge, G., Cottens, S., Weitz-Schmidt, G., and Hommel, U. (1999) *J. Mol. Biol.* **292**, 1-9.
35. Weitz-Schmidt, G., Welzenbach, K., Brinkmann, V., Kamata, T., Kallen, J., Bruns, C., Cottens, S., Takada, Y., and Hommel, U. (2001) *Nat. Med.* **7**, 687-692.
36. Ajuebor, M. N., A. M. Das, L. Virag, R. J. Flower, C. Szabo, and M. Perretti. (1999). *J. Immunol.* **162**, 1685.

Table I: Pepspot analysis of peptides derived from MMPs and $\alpha_M\beta_2$ integrin ligands (SEQ ID NOS: 31-100, 174, 12, 5, 6, 101, 102, 103, and 104 respectively)

Protein	Peptide	OD/mm ²	Binding positivity
MMP-1	DAHFDEDERWTNNFR	1792 +	
	DAHFDEAARWTNNFR	1417 -	
MMP-2	DSHFDDDELWTLGEG	4687 +++	
	DSHFDDAALWTLGEG	3334 ++	
MMP-3	DAHFDDDEQWTKDTT	4993 +++	
	DAHFDDAAQWTKDTT	2188 +	
MMP-7	DAHFDEDERWTDGSS	4043 ++	
	DAHFDEAARWTDGSS	2065 +	
MMP-8	DAHFDAEETWTNTSA	4295 ++	
	DAHFDAAAATWTNTSA	1258 -	
MMP-9	DAHFDDDELWSLGKG	2010 +	
	DAHFDDAALWSLGKG	476 -	
MMP-13	DAHFDDDETWTSSSK	4324 ++	
	DAHFDDAATWTSSSK	1732 +	
MMP-14	DTHFDASAEPWTVRNE	1264 -	
	DTHFDASAAPWTVRNE	1198 -	
MMP-1	SGDVQLDDIDGIQAI	484 -	
	SGDVQLAAIDGIQAI	441 -	
MMP-3	RFLRLSQDDINGIQSL	810 -	
	RFLRLSQAAINGIQSL	1541 +	
MMP-8	NYSLPQDDIDGIQAI	3348 ++	
	NYSLPQAIDGIQAI	505 +	
MMP-13	HFMLPDDDVQGIQSL	542 -	
	HFMLPDAAVQGIQSL	384 -	
MMP-14	NFVLPDDDRRGIQQL	518 -	
	NFVLPDAARRGIQQL	609 -	
Fibronectin	HEATCYDDGKTYHVG	1271 -	
	HEATCYAAGKTYHVG	596 -	
ICAM-3	LNATESDDGRSFFCS	369 -	
	LNATESAAGRSFFCS	277 -	
Complement factor H	EEMHCSDDGFWSKEK	2972 +	
	EEMHCSAAGFWSKEK	321 -	
TSP-1	WPSDSADDGWSPWSE	4655 +++	
	WPSDSAAAGWSPWSE	1543 +	
NIF	DPVCIPDDGVCFIGS	221 -	
	DPVCIPAAGVCFIGS	114 -	
ICAM-2	NSTADREDGHNFSC	75 -	
	NSTADRAAGHRNFSC	73 -	
Fibronectin	NVYQISEEDGEQSLIL	1260 -	
	NVYQISAAGEQSLIL	453 -	
Fibronectin	VTYSSPEDGIHELP	301 -	
	VTYSSPAAGIHELP	212 -	
Cyr61	KMRFRCEDGETFSKN	317 -	
	KMRFRCAAGETFSKN	386 -	

Myeloperoxidase	WLPAEYEDGFSLPYG	4603	+++
Catalase	WLPAEYAAGFSLPYG	924	-
	AVKFYTEDGNWDLVG	5045	+++
	AVKFYTAAGNWDLVG	690	-
Fibrinogen alpha	KEVVTSEDGSDCPEA	124	-
	KEVVTSAAGSDCPEA	225	-
Fibrinogen beta	RKQCSKEDGGGWYN	483	-
	RKQCSKAAGGGWWY	325	-
Fibrinogen alpha	N GFGSLNDEGEGEFWL	732	-
	GFGSLNAAGEGEFWL	397	-
GP1b	GCPTLGDEGDTDLYD	544	-
	GCPTLGAAAGDTDLYD	238	-
ICAM-1	VSVTAEDEGTQRLTC	51	-
	VSVTAEAAGTQRLTC	68	-
Factor X	DRNTEQEEGGGEAVHE	379	-
	DRNTEQAAGGEAVHE	236	-
E-selectin	TCTFDCEEGFELMGA	309	-
	TCTFDCAAGFELMGA	70	-
E-selectin	SCNFTCEEGLQGP	138	-
	SCNFTCAAGFMLQGP	54	-
E-selectin	SCAFSCEEGLHGS	236	-
	SCAFSCAAGFELHGS	74	-
Fibronectin	TFHKRHEEGHMLNCT	39	-
	TFHKRHAAGHMLNCT	12	-
Fibronectin	VEYELSEEGDEPQYL	3676	++
	VEYELSAAGDEPQYL	2682	+

Range: 0-1499 OD/mm² = -, 1500-2999 = +, 3000-4499 = ++,
>4500 = +++

	Peptide	No competitor	Soluble DDGW
DDGW peptide	ADGACILWMDDGWC	10859	0
iC3b	GAAG		
	ARSNLDDEDIIAEENI	13265	409
iC3b	ARSNLDAIIIAEENI	0	0
	EDIIAEENIVSRSEF	0	0
iC3b	EDIIAAANIVSRSEF	0	0
	EGVQKEDIPPADLSD	0	0
iC3b	EGVQKAAIPPADLSD	0	0